

1 **Control of N₂ fixation and NH₃ excretion in *Azorhizobium caulinodans* ORS571**

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11 **Abstract**

12 Due to the costly energy demands of N₂ fixation, diazotrophic bacteria have evolved complex
13 regulatory networks that permit expression of the N₂-fixing catalyst nitrogenase only under conditions
14 of N starvation, whereas the same condition stimulates upregulation of high-affinity NH₃ assimilation
15 by glutamine synthetase (GS), preventing excess release of excess NH₃ for plants. Diazotrophic
16 bacteria can be engineered to excrete NH₃ by interference with GS, however control is required to
17 minimise growth penalties and prevent unintended provision of NH₃ to non-target plants. Here, we
18 attempted two strategies to control GS regulation and NH₃ excretion in our model cereal symbiont
19 *Azorhizobium caulinodans* AcLP, a derivative of ORS571. We first attempted to recapitulate previous
20 work where mutation of both P_{II} homologues *glnB* and *glnK* stimulated GS shutdown but found that
21 one of these genes was essential for growth. Secondly, we expressed unidirectional
22 adenylyltransferases (uATs) in a Δ *glnE* mutant of AcLP which permitted strong GS shutdown and
23 excretion of NH₃ derived from N₂ fixation and completely alleviated negative feedback regulation on
24 nitrogenase expression. We placed a *uAT* allele under control of the NifA-dependent promoter *PnifH*,
25 permitting GS shutdown and NH₃ excretion specifically under microaerobic conditions, the same cue
26 that initiates N₂ fixation, then deleted *nifA* and transferred a rhizopine-inducible *nifA*_{L94Q/D95Q}-*rpoN*
27 controller plasmid into this strain, permitting coupled rhizopine-dependent activation of N₂ fixation
28 with NH₃ excretion. In future, this highly sophisticated and multi-layered control circuitry could be
29 used to activate N₂ fixation and NH₃ excretion specifically by AcLP colonising transgenic rhizopine
30 producing cereals, targeting delivery of fixed N to the crop, and preventing interaction with non-target
31 plants.

32

33 **Author Summary**

34 Inoculation of cereal crops with associative “diazotrophic” bacteria that convert atmospheric N₂ to
35 NH₃ could be used to sustainably improve delivery of nitrogen in agriculture. However, due to the
36 costly energy demands of N₂ fixation, natural diazotrophic bacteria have evolved to conserve energy
37 by preventing excess production of NH₃ and release to the plants. Diazotrophs can be engineered for
38 excess NH₃ production and release, however genetic control is required to minimise growth penalties
39 and prevent unintended provision of NH₃ to non-target weed species. Here, we engineer control of N₂
40 fixation and NH₃ excretion in response to the signalling molecule rhizopine which is produced by
41 transgenic barley. This control could be used to establish plant host-specific activation of N₂ fixation
42 and NH₃ release following root colonisation in the field, minimising bacterial energy requirements in
43 the bulk soil and preventing provision of NH₃ to non-target plants.

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46 Introduction

47 Nitrogen (N) is an essential constituent of all biological organisms, but metabolically accessible forms
48 are scarce in most environments, restricting biomass production. In agriculture, productivity of cereal
49 crops, which are a staple of human dietary requirements, requires large-scale supplementation with
50 synthetic N fertilisers to meet global food security requirements (1). However, synthesis and
51 excessive application of N fertilisers has a large energy cost, causes CO₂ release and results in loss of
52 reduced N to the environment, which has doubled reactive N in the atmosphere and polluted
53 waterways causing eutrophication and O₂-depleted dead zones (2). In contrast, N fertilisers are largely
54 unaffordable to small-hold farmers in developing countries such as those in Sub-Saharan Africa (3),
55 restricting yields to a fraction of their maximum potential (4). Inoculation of cereals with root-
56 associative diazotrophic bacteria that convert atmospheric N₂ gas to NH₃ through the action of O₂-
57 labile nitrogenase represents an affordable and sustainable alternative to the use of N fertilisers in
58 agriculture (5-7). Although associative diazotrophs have been estimated to fix up to 70 kg N ha⁻¹ year⁻¹
59 in agricultural systems (8), responses to inoculation are typically inconsistent due to sub-optimal
60 competitiveness for root colonisation and persistence in soil (9-12). Furthermore, due to the costly
61 energy demands of N₂ fixation, which consumes at least 16 mol ATP per mol N₂ fixed *in vitro*,
62 bacteria have evolved complex regulatory networks that permit expression and activity of the N₂-
63 fixing catalyst nitrogenase only under conditions of N starvation, whereas the same condition
64 stimulates upregulation of high-affinity NH₃ assimilation by glutamine synthetase (*glnA*, GS),
65 preventing excess release of excess NH₃ for plants (13, 14).

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67 Associative diazotrophic bacteria can be engineered for excess production and excretion of NH₃ by
68 several strategies (13, 15, 16). For example, in *Azotobacter vinelandii*, insertional inactivation of *nifL*,
69 which encodes an O₂ as well as N and C sensing anti-activator of the nitrogenase master regulator
70 NifA, drives constitutive nitrogenase activity resulting in excretion of NH₃ from the cell (17-20). The
71 same effect was achieved by expressing mutant *nifA* alleles that are resistant to inhibition by NifL (18,
72 21, 22). While excess NH₃ production itself is likely to activate regulatory feedback mechanisms

73 reducing GS biosynthetic activity and NH₃ assimilation (15), mutating *glnA* (23-27) or genes involved
74 in GS regulation may also be required to inhibit NH₃ assimilation more strongly and favour NH₃
75 excretion (28, 29).

76

77 Bacterial GS belongs to the “class I” type enzymes comprised of 12 identical subunits which are each
78 adenylylated or de-adenylylated by a bidirectional adenylyl transferase (AT, encoded by *glnE*) at the
79 Tyr₃₉₇ residue, with the fully de-adenylylated GS form being biosynthetically active and vice versa
80 (30). Directionality of the ATase reaction is regulated by the post-translational modification state of P_{II}
81 signal transduction proteins (31). The activity of P_{II} proteins is regulated by
82 uridylylation/deuridylylation by the bidirectional uridylyltransferase (UT) GlnD which represents the
83 most basal regulator in the cascade and can directly sense N status of the cell (32). GlnD uridylylates
84 P_{II} under conditions of N-starvation and the resulting P_{II}-UMP ultimately triggers dephosphorylation
85 of ATase and hence deadenylylation and activation of GS (33). In *Azorhizobium caulinodans* (*Ac*),
86 insertional inactivation of both P_{II} homologues *glnB* and *glnK* produced a mutant that was unable to
87 activate GS by deadenylylation, driving NH₃-insensitive N₂ fixation and excretion of NH₃ into the
88 growth media (28). Critically, this engineering strategy does not appear to be universally applicable as
89 P_{II} is essential for NifA and nitrogenase activity in some bacteria (34, 35), whereas it is essential for
90 growth in others (36, 37). In a Δ *glnE* ATase mutant of *Azospirillum brasilense*, complementation with
91 unidirectional adenylyltransferase (uAT) alleles that encoded only the C-terminal adenylylation domain
92 (31) drove strong adenylylation of GS resulting in excretion of NH₃ into the growth media (29). This
93 strategy likely represents a more universally applicable approach for engineering NH₃ excretion in
94 diazotrophs because the ATase is highly conserved, has a specific function, and can be readily mutated
95 across diverse diazotrophic bacterial taxa (15, 38-40), albeit the mutation appears to be lethal in the
96 heterotroph *Mycobacterium tuberculosis* (41, 42).

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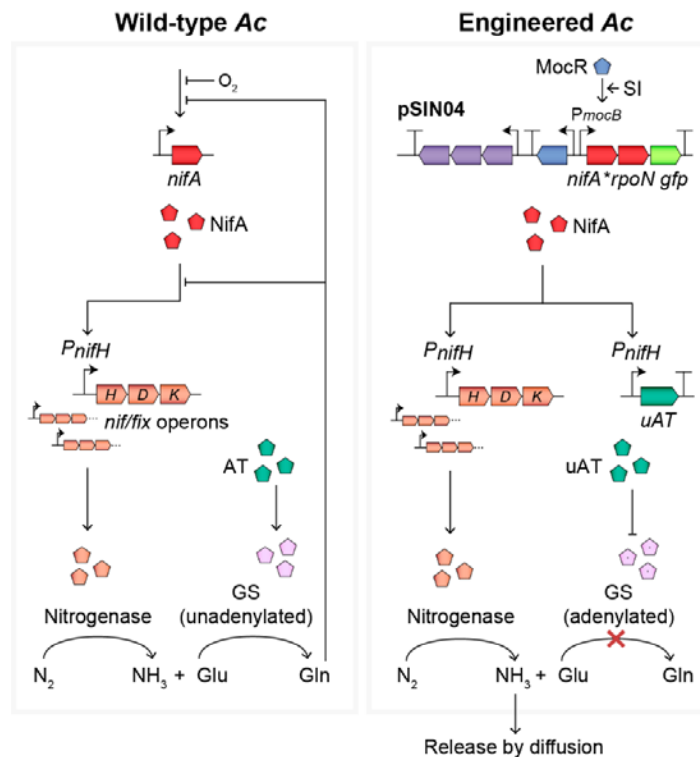
98 From an agricultural perspective, there are three major caveats of engineering diazotrophic bacteria
99 for excessive production and excretion of NH₃; i) uncontrolled *nifA* and(or) nitrogenase expression
100 has a severe energy burden on the cell that could abolish competitiveness for root colonisation; ii)

101 interference with GS activity typically renders strains auxotrophic for the essential amino acid
102 glutamine, which could further reduce competitiveness; and iii) NH₃ excreting bacteria have potential
103 to supply NH₃ to non-target weed species following promiscuous colonisation in the field. Therefore,
104 establishing control of N₂ fixation and NH₃ excretion will be crucial for the optimisation of strains as
105 agricultural inoculants. Control of NH₃ excretion has already been achieved in *A. vinelandii* by
106 establishing IPTG-dependent expression of *glnA* (27), and in *A. brasilense* by establishing anhydro-
107 tetracycline inducible expression of uATs (29, 43). However, use of plant-derived signals to control
108 N₂-fixation and NH₃ excretion would be far more applicable in the environment and could impart
109 partner-specificity to target delivery of fixed N to crops and prevent interactions with non-target host
110 plants following promiscuous colonisation (44, 45).

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112 We previously developed synthetic rhizopine signalling between barley and the model endophyte
113 *Azorhizobium caulinodans* AcLP that stimulates transcriptional activation of the mutant nitrogenase
114 master regulator *nifA*_{L94Q/D95Q}, which partially escapes nitrogen regulation, and when paired with the
115 sigma factor RpoN drives N₂ fixation in bacteria colonising rhizopine producing (*RhiP*) barley roots
116 (44, 46, 47). Here, we demonstrate that wild-type and engineered *Ac* strains do not release fixed N as
117 NH₃ into the growth media when cultured under N₂-fixing conditions and therefore sought to engineer
118 this trait by interfering with high-affinity NH₃ assimilation catalysed by GS. In our attempts to
119 recapitulate NH₃ excreting *glnB glnK* double mutants of AcLP (28), we found that deletion of both P_{II}
120 homologues was only possible when second copy of *glnB* was first integrated into the chromosome
121 suggesting one of the P_{II} homologues were essential for growth. GS and nitrogenase activity in the
122 resulting strain exhibited minimal variation from that of the wild-type, but nevertheless the strain
123 excreted low levels of NH₃ into the growth media. To optimise rates of NH₃ excretion, we utilised a
124 second engineering strategy where a *AcLPΔglnE* mutant was complemented with uATs. In
125 congruency with similar experiments performed in *A. brasilense* (29), uAT expression drove strong
126 shutdown of GS, but also completely alleviated negative feedback inhibition of nitrogenase by NH₃
127 and stimulated NH₃ excretion. By placing uAT expression under control of NifA, we established
128 control of these traits by the same cues that initiate N₂ fixation, then transferred rhizopine control of

129 *nifA_{L94Q/D95Q}rpoN* into this strain linking activation of N₂-fixation and NH₃ excretion (Fig 1). This
 130 highly sophisticated control circuitry represents a significant milestone in the development of a
 131 “synthetic symbiosis” where N₂ fixation and NH₃ excretion could be activated in bacteria specifically
 132 colonising target cereals, targeting delivery of N to the crops while avoiding potential interactions
 133 with non-target plants.
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137 **Figure 1. Model for rhizopine control of N₂ fixation and NH₃ excretion in *Ac*.** In the wild-type
 138 bacterium, NifA is activated under N₂-fixing (N-free microaerobic) conditions leading to transcription
 139 of nitrogenase (*nif* and *fix*) genes and subsequently N₂ fixation. Under the same conditions, P_{II}-UMP
 140 stimulates the adenylyltransferase (AT) to activate glutamine synthetase (GS) by deadenylylation. GS
 141 catalyses assimilation of NH₃ via the conversion of glutamate (Glu) to glutamine (Gln), which feeds
 142 back to repress *nif* expression and NifA activity, preventing excess production and release of NH₃
 143 from the cell. In our engineered strain *Ac*PU-R22 carrying pSIN04, *nifA_{L94Q/F95Q}* is expressed by

144 addition of rhizopine to the culture, driving nitrogenase expression and N₂ fixation. Additionally, NifA
145 induces transcription of the unidirectional adenylyltransferase (uAT) which drives shutdown of GS by
146 adenylylation, preventing NH₃ assimilation. Because shutdown of GS prevents glutamine
147 biosynthesis, repression on *nif* gene expression and NifA activity is alleviated, resulting in excess
148 production of NH₃ and release from the cell.

149

150 **Results**

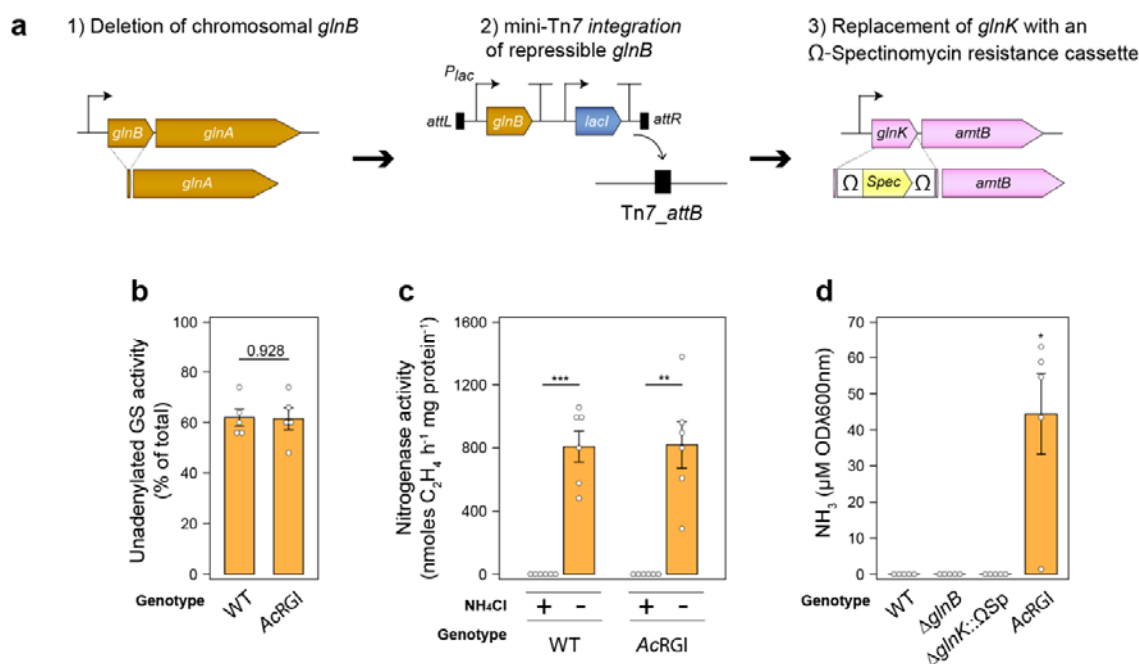
151 **Deletion or strong repression of the *P_{II}* genes is lethal**

152 It was previously demonstrated that insertional inactivation of the *Ac P_{II}* genes *glnB* and *glnK*
153 stimulates shutdown of GS by adenylylation and alleviates negative feedback inhibition of
154 nitrogenase by the product NH₃, preventing NH₃ assimilation and favouring excretion into the growth
155 media (28). We attempted to recapitulate these experiments in *AcLP*, a derivative of *Ac* harbouring a
156 mini-Tn7 *attB* integration site stably recombined into its chromosome, by constructing a markerless
157 deletion of *glnB* and replacing *glnK* with an omega (Ω)-spectinomycin resistance (Sp) cassette.
158 Although the single Δ *glnB* and Δ *glnK*:: Ω Sp mutations were readily acquired, we were unable to
159 acquire the double mutant by introduction of the Δ *glnK*:: Ω Sp mutation into *AcLP* Δ *glnB* when
160 selection was performed on rich or minimal media supplemented with glutamine as a sole N source,
161 suggesting the resulting phenotype was lethal. To explore this notion further, we integrated into the
162 chromosome of *AcLP* Δ *glnB* a construct encoding *glnB* with the strong ribosome binding site (RBS)
163 RStd expressed from the IPTG de-repressible promoter *Plac* (Fig 2a) and were subsequently able to
164 acquire the Δ *glnB* Δ *glnK*:: Ω Sp double mutation when selection was performed on rich media in the
165 absence of IPTG, confirming that one of the *P_{II}* proteins was essential for growth.

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Figure 2. Strong repression of *glnB* in a *glnK* mutant has minimal effect on glutamine synthetase

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and nitrogenase activity but drives low-level NH₃ excretion. (a) Strategy for generating strain

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AcRGI with the double Δ *glnB* and Δ *glnK*:: Ω Sp mutation following integration of an IPTG-

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derepressible *glnB* gene into the chromosome of *AcLP*. **(b)** Activity of the unadenylylated (active)

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form of GS in $n = 5$ wild-type (WT) or *AcRGI* cultures incubated for 24-h as determined by γ -

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glutamyl transferase assays in the presence or absence of 60 mM MgCl₂ (see S2 Fig for total activity).

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(c) Nitrogenase activity measured by acetylene reduction in $n = 6$ cultures between 3-h – 21-h **(d)**

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Spectrophotometric determination of NH₃ in media of $n = 5$ cultures grown for 24-h. Cultures for all

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assays were grown in N₂-fixing conditions (N-free UMS media with 3% O₂ in the headspace). Error

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bars represent one SEM. Independent two-tailed students t-tests were used to compare means. Exact

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P-values are provided where $P > 0.05$. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$. The wild-type *AcLP* was

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used as a reference group for comparison of means in panel (e).

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We next sought to test whether reduced translation of the introduced *glnB* module would stimulate GS

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shutdown and NH₃ excretion by tuning the ribosome binding site (RBS). Seven synthetic RBS' were

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experimentally demonstrated to produce translation rates spanning two to three orders of magnitude

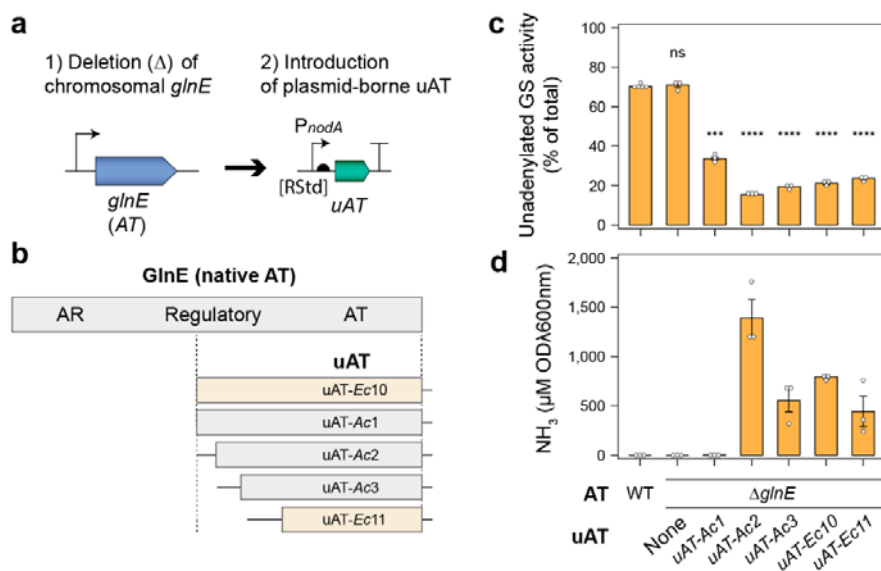
186 (S1 Fig), but only when *glnB* fused to the strongest RBS “RStd” and integrated into the *AcLPΔglnB*
187 chromosome were we able to subsequently isolate the $\Delta glnK::\Omega Sp$ mutation (hereby termed strain
188 *AcRGI*), suggesting that *glnB* had been repressed as much as was tolerable. We assessed total GS
189 specific activity and that of the unadenylylated active enzyme in *AcRGI* by performing γ -glutamyl
190 transferase assays on whole cells in the presence or absence of 60 mM $MgCl_2$ which specifically
191 inhibits the adenylylated enzyme (48), and found that mutant exhibited higher total GS activity
192 compared to the wild-type (S2a Fig), presumably due to elevated *glnA* expression (S2b Fig) as is
193 typical of *glnB* mutants (25, 28), whereas the adenylylation state of GS (depicted here as percentage
194 of unadenylylated GS activity) was unchanged (Fig 2b). We also found that the specific nitrogenase
195 activity of strain *AcRGI* was no different from that of the wild-type, being repressed by
196 supplementation of 10 mM NH_3Cl into the growth media (Fig 2c). Spectrophotometric quantification
197 of NH_3 was next performed using the indophenol method (49) on the strains grown for 24-h under N_2 -
198 fixing conditions (here defined as N-free UMS with O_2 in the headspace adjusted to 3%). No NH_3 was
199 detected in the wild-type or $\Delta glnB$ and $\Delta glnK::\Omega Sp$ single mutants, whereas we detected trace
200 amounts of NH_3 in the growth media of strain *AcRGI* (Fig 2d). Given that construction of the
201 *AcLPΔglnB ΔglnK::ΩSp* double mutant was lethal, and strong repression of *glnB* had minimal effect
202 on GS and nitrogenase regulation permitting only low-level NH_3 excretion, we concluded that these
203 strategies were inadequate to establish control of NH_3 excretion in *AcLP* and opted to pursue an
204 alternative strategy.

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206 **uAT expression drives GS inactivation and NH_3 excretion**

207 In a $\Delta glnE$ mutant of *A. brasilense*, controlled expression of a N-terminal truncated AT consisting of
208 only the adenylylation domain results in unidirectional activity driving strong inactivation of GS by
209 adenylylation and excretion of NH_3 into the growth media (29). We recapitulated these experiments in
210 a $\Delta glnE$ mutant of *AcLP* by using the *Sinorhizobium meliloti* derived *PnodA* promoter (S3 Fig), to
211 drive expression of a series of truncated uATs derived from *Ac* or those previously described for *E.*
212 *coli* (Fig 3a-b) (29). We assessed GS specific activity and that of the unadenylylated enzyme using γ -

213 glutamyl transferase assays on cells grown in N₂-fixing conditions for 3-h and confirmed that leaky
 214 non-induced uAT expression stimulated GS adenylation (Fig 3c), while having minimal effect on
 215 total GS specific activity relative to wild-type bacteria (S4 Fig). The strains also excreted between
 216 0.1-1.5mM of NH₃ after 24-h incubation in N₂-fixing conditions, whereas the wild-type and $\Delta glnE$
 217 mutant did not excrete detectable levels of NH₃ (Fig 3d). Interestingly, we found that NH₃ excretion
 218 was sub-optimal when the *PnodA* promoter controlling *uAT* expression was induced with saturating
 219 levels (5 μ M) of naringenin (S5 Fig), suggesting that *uAT* overexpression is metabolically detrimental,
 220 as was observed in *A. brasilense* (43). This indicated that more finely tuned *uAT* expression would be
 221 critical to achieve stringent control of GS adenylation in *AcLP*.
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223
 224 **Figure 3. uAT expression drives GS adenylation and NH₃ excretion in a $\Delta glnE$ background.** (a)
 225 Strategy for complementation of the $\Delta glnE$ mutation with naringenin-inducible unidirectional
 226 adenylyltransferases (uAT) expressed from low-copy parent plasmid pOPS1536. (b) A series of
 227 truncated uAT proteins were used in this study. The *uAT-Ec10* and *uAT-Ec11* alleles are derived from
 228 *E. coli* were described previously (29), whereas *uAT-Ac* alleles are derived from *AcLP*. The nucleotide
 229 sequences for these alleles are provided in S1 File. (c) Activity of the unadenylylated (active) form of
 230 GS in $n = 5$ for *AcLP* (wild-type, WT) or $n = 3$ cultures incubated for 3-h in N₂-fixing conditions (N-
 231 free UMS media with 3% O₂ in the headspace) without the inducer naringenin as determined by γ -

232 glutamyl transferase assays in the presence or absence of 60 mM MgCl₂ (see S4 Fig for total activity).
233 (d) Spectrophotometric determination of NH₃ in media of $n = 3$ cultures grown for 24-h in N₂-fixing
234 conditions. Error bars represent one SEM. Independent two-tailed students t-tests with the Bonferroni-
235 holm adjustment were used to compare means using the wild-type *AcLP* as a reference group. Not
236 significant (ns $P > 0.05$), *** $P < 0.001$, **** $P < 0.0001$.

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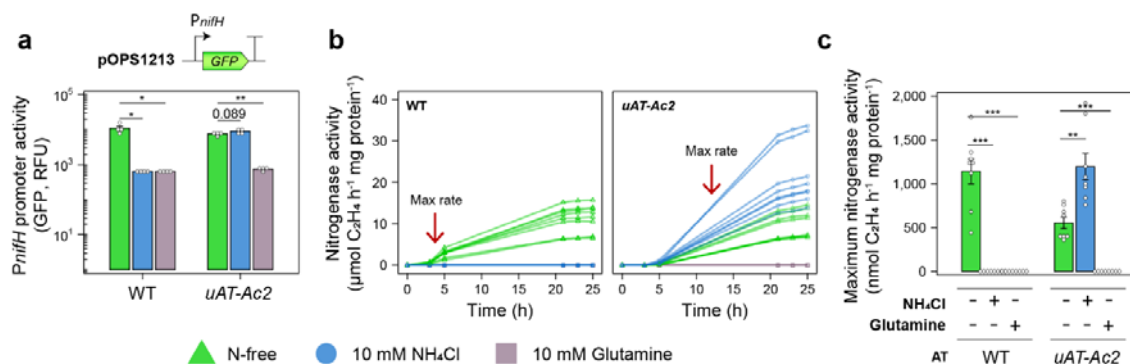
238 **Shutdown of glutamine biosynthesis alleviates negative feedback** 239 **on nitrogenase**

240 Expression of *uAT* restricts glutamine production via the high affinity GS-dependent NH₃ assimilation
241 pathway, providing us with a unique opportunity to tease apart the effects of NH₃ and glutamine on
242 the nitrogenase (*nif*) gene expression. We postulated that NH₃ must first be converted into glutamine
243 to mediate repression of *nif* genes and tested this hypothesis first by examining expression of *PnifH*
244 fused to *GFP* on plasmid pOPS1213 in wild-type bacteria and in *AcLPΔglnE* expressing the *uAT-Ac2*
245 allele from the non-induced *PnodA* promoter on a second plasmid. As expected, *PnifH::GFP* activity
246 in both strains grown under microaerobic conditions (3% O₂ in the headspace) was strongly repressed
247 by supplementation with 10 mM glutamine however, while *PnifH::GFP* was repressed in the wild-
248 type by supplementation with 10 mM NH₄Cl, *PnifH::GFP* expression was not repressed by NH₄Cl in
249 *AcLPΔglnE* expressing *uAT-Ac2* (Fig 4a). We observed a similar pattern when nitrogenase activity
250 was assessed by ARAs (Fig 4b-c), indicating that NH₃ itself has no effect on negative feedback
251 regulation of *nif* genes but must be converted into glutamine or potentially other amino acids to
252 facilitate repression. Engineering NH₃ excreting bacteria by targeted GS shutdown therefore has two
253 advantages; i) alleviating negative feedback regulation of *nif* genes and ii) preventing NH₃
254 assimilation to favour release.

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260 **Figure 4. uAT expression abolishes negative feedback regulation on nitrogenase.** (a) A
 261 *PnifH::GFP* reporter carried on plasmid pOPS1213 was mobilised into the wild-type (WT) AcLP and
 262 AcLP Δ *glnE* expressing *uAT-Ac2* on a second low-copy plasmid and induction was measured in $n = 4$
 263 cultures grown for 24-h under the conditions indicated. Relative fluorescence units (RFU) are defined
 264 here as GFP fluorescence / OD_{600nm} (b) Nitrogenase activity was measured by acetylene reduction
 265 in $n = 8$ cultures grown under N_2 -fixing conditions (N-free UMS media with 3% O_2 in the headspace)
 266 and (c) the maximum rates are presented. Error bars represent one SEM. Independent two-tailed
 267 students t-tests with Bonferroni-holm adjustment were used to compare means. Exact P-values are
 268 provided where $P > 0.05$. ** $P < 0.01$, *** $P < 0.001$. WT bacteria grown at 3% O_2 in N-free conditions
 269 was used as a reference group for statistical comparisons in panel (a).

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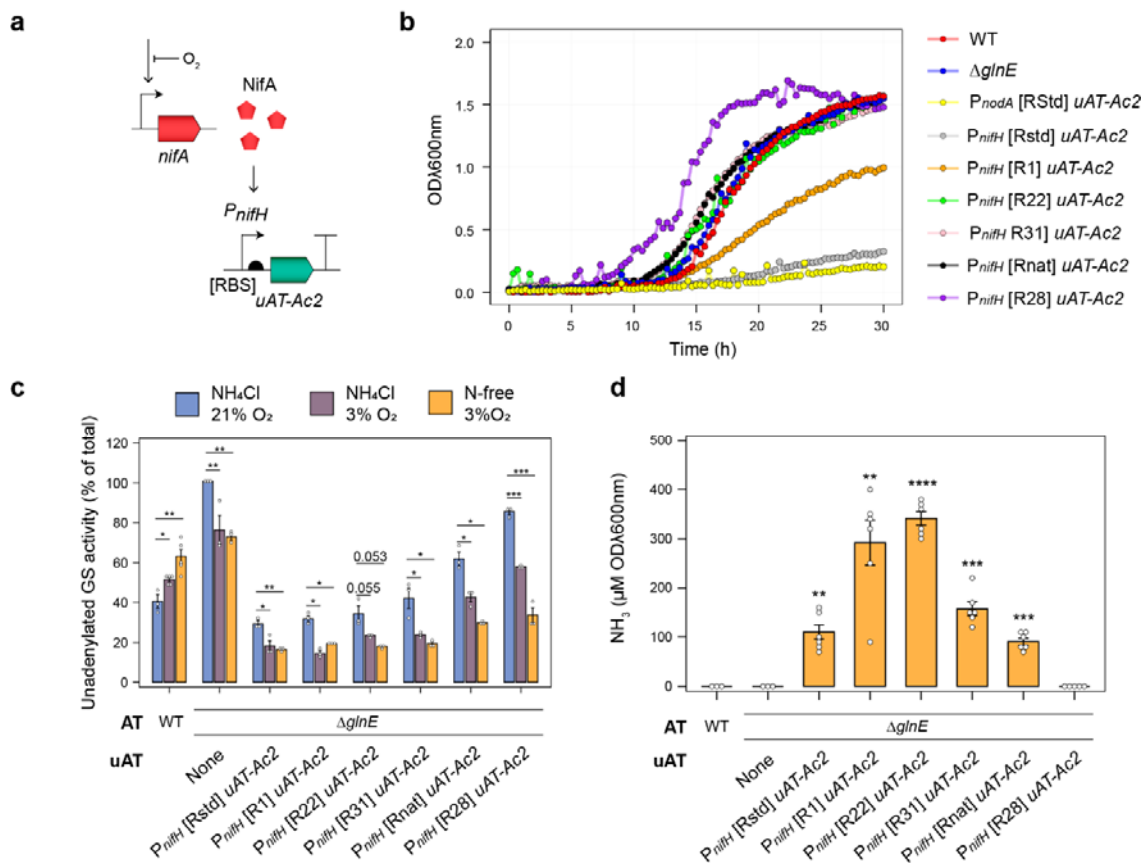
271 **NifA control of uAT expression**

272 As a direct consequence of engineering NH_3 excretion through GS interference, bacteria typically
 273 become auxotrophic for glutamine. While this may not be non-problematic for cultures grown *in vitro*
 274 under gnotobiotic conditions, glutamine auxotrophs in the field would be unable to compete or persist
 275 in the soil and rhizosphere. In rhizobia-legume symbioses, rhizobia only restrict NH_3 assimilation
 276 after infecting the low-oxygen environment of the nodule and differentiating into an N_2 fixing
 277 bacteroid (50, 51), allowing them to maintain competitiveness during their free-living state in the soil.
 278 To mimic this regulation, we fused the *uAT-Ac2* allele to native or synthetic RBSs and placed these

279 under control of the NifA-inducible *PnifH* promoter on mini-Tn7 delivery plasmids, then integrated
280 these into the chromosome of *AcLPΔglnE*, creating strains *AcPU-RStd*, *AcPU-R1*, *AcPU-R22*, *AcPU-*
281 *R31*, *AcPU-Rnat* and *AcPU-R28* (Fig 5a). When grown under aerobic (21% O₂) conditions in the
282 presence of 10 mM NH₄Cl, growth of *AcLPΔglnE* expressing the *uAT-Ac2* allele from the non-
283 induced *PnodA* promoter was almost entirely abolished compared to where glutamine was provided as
284 a source of N (Fig 5b and S6 Fig). In contrast, the growth characteristics of strains expressing *uAT-*
285 *Ac2* from the *PnifH* promoter were reminiscent of the wild-type *Ac*, except for strains where *uAT-Ac2*
286 was fused to the strongest RBS' RStd or R1, which increased mean generation times (MGT) but did
287 not affect the total biomass at stationary phase (Fig 5b and S6 Fig). We next assessed GS regulation
288 by γ -glutamyltransferase assays and confirmed that under aerobic conditions in the presence of 10
289 mM NH₄Cl, the percentage of active deadenylylated GS activity in strains *AcPU-R1*, *AcPU-R22*, and
290 *AcPU-R3* closely resembled that of the wild-type, suggesting that NH₃ assimilation was functional.
291 When grown under microaerobic conditions (3% O₂) in the presence or absence of 10 mM NH₄Cl, GS
292 in wild-type *AcLP* was activated by de-adenylylation, whereas GS in all *AcLPΔglnE* strains
293 expressing *uAT-Ac2* from the *PnifH* promoter became more heavily inactivated by adenylylation
294 under the same conditions (Fig 5c), with the percentage unadenylated GS activity correlating
295 negatively with the strength of RBS fused to *uAT-Ac2*. We finally performed NH₃ excretion assays on
296 the engineered strains and found that each excreted NH₃ into the growth media after 24-h, except for
297 where *uAT-Ac2* was fused to the weakest RBS [R28] (Fig 5d). Overall, the data suggested that by
298 expressing *uATs* from the *PnifH* promoter, GS shutdown can be controlled in response to the same
299 cues that initiates N₂ fixation.

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303 **Figure 5. Coupled activation of N₂ fixation and GS adenylylation via NifA-dependent expression**

304 **of uAT. (a)** Strategy for complementation of the $\Delta glnE$ mutation with NifA-inducible unidirectional

305 adenylyltransferases (uAT) integrated into the chromosome using mini-Tn7. **(b)** Growth of control

306 strains and those expressing uATs in UMS media supplemented with 20 mM succinate and either 10

307 mM glutamine or 10 mM NH₃Cl under aerobic conditions. See S6 Fig for full growth statistics. **(c)**

308 Activity of the unadenylylated (active) form of GS in $n = 5$ for wild-type (WT) AcLP or $n = 3$ cultures

309 incubated for 24-h in as determined by γ -glutamyl transferase assays in the presence or absence of 60

310 mM MgCl₂. **(d)** Spectrophotometric determination of NH₃ in media of $n = 3$ WT and $\Delta glnE$ or $n = 5$

311 cultures grown for 24-h in N₂-fixing conditions (N-free UMS media with 3% O₂ in the headspace).

312 Error bars represent one SEM. Independent two-tailed students t-tests with the Bonferroni-holm

313 adjustment were used to compare means using the wild-type AcLP as a reference group. Exact P-

314 values are provided where $P > 0.05$. ** $P < 0.01$, *** $P < 0.001$. WT bacteria were used as a reference

315 group for statistical comparisons in panels (a) and (b).

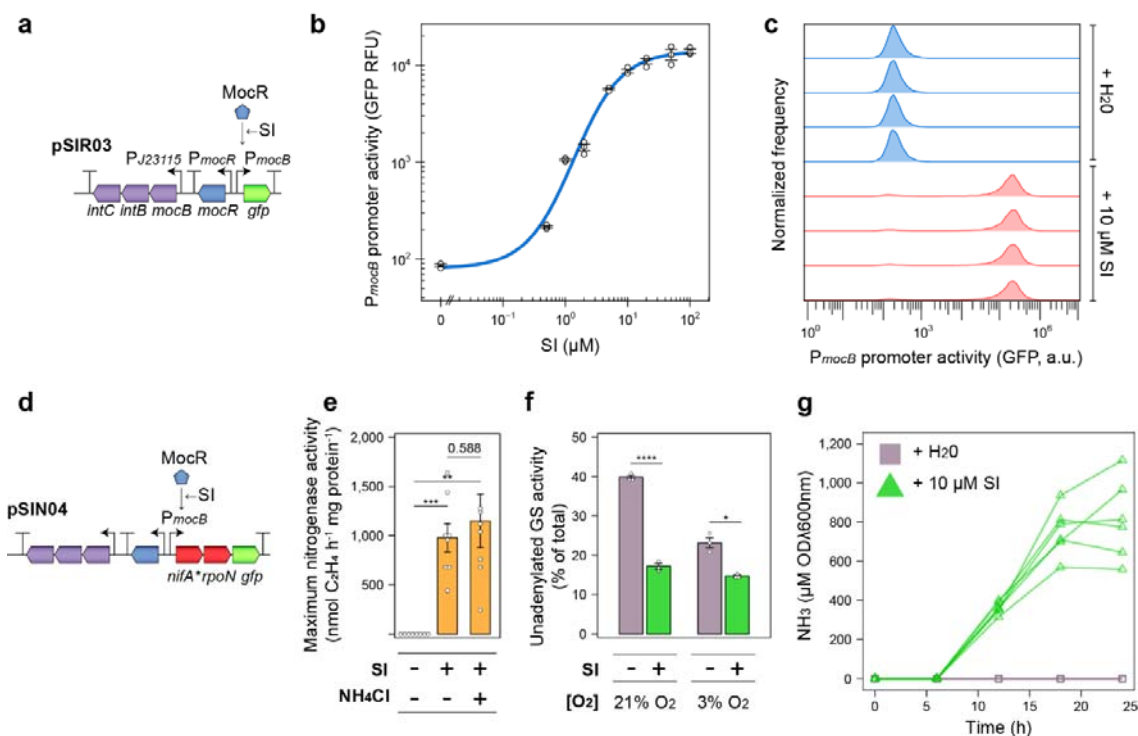
316

317 **Rhizopine-dependent control of N₂ fixation, GS adenylation and**

318 **NH₃ excretion**

319 While NifA-dependent expression of nitrogenase and *uAT-Ac2* in *AcΔglnE* drives N₂ fixation and GS
320 inactivation leading to NH₃ excretion, the lack of plant host-specific signalling to drive these
321 processes could permit bacteria to supply NH₃ to target crops and non-target weed species alike. We
322 previously used synthetic rhizopine signalling to establish control of a mutant *nifA* allele (encoding
323 NifA_{L94Q/D95Q}) and *rpoN* in *AcLPΔnifA* carrying plasmid pSIN02, which drove partially NH₃-resistant
324 activation of nitrogenase activity specifically by bacteria occupying the roots of transgenic *RhiP*
325 barley (46). We performed NH₃ excretion assays on *AcLPΔnifA* carrying pSIN02 and found that this
326 strain did not secrete NH₃ into the growth media (S7 Fig). Thus, we opted to establish rhizopine
327 control of the *nifA_{L94Q/D95Q}-rpoN* operon in our strain *AcPU-R22* where *uAT-Ac2* expression placed
328 under control by NifA. We first tested in *AcLP*, induction of a new rhizopine receiver plasmid pSIR03
329 which was derived from the high-copy rhizopine receiver pSIR03 but carried an RK2 replicon for
330 low-copy maintenance. Using *GFP* induction assays, we demonstrated that pSIR03 (Fig 6a) has a
331 dynamic range of 162-fold in response to the rhizopine *scyllo*-inosamine (SI) and was induced in
332 93.08 ± [SEM] 0.32% of cells in populations when 10 μM SI was supplemented *in vitro* (Fig 6b-c &
333 Table S1). We deleted the native *nifA* gene from strain *AcPU-R22* and introduced a rhizopine
334 *nifA_{L94Q/D95Q}-rpoN* controller plasmid pSIN04 which was derived from pSIR03 (Fig 6d). Expression of
335 *nifA_{L94Q/D95Q}-rpoN* under microaerobic conditions by addition of 10 μM SI into the media resulted in
336 tightly controlled activation of nitrogenase that was unimpeded by addition of 10 mM of NH₃ (Fig
337 6e). Moreover, GS was strongly adenylylated by addition of 10 SI to the media in both aerobic and
338 microaerobic conditions (Fig 6f). Because NifA in many diazotrophs is inactivated when cells are
339 grown at 21% O₂ (52), we subsequently tested O₂ tolerance of our NifA_{L94/D95Q} mutant protein by
340 inducing expression of *nifA_{L94Q/D95Q}-rpoN* in *AcLPΔnifA* carrying pSIN03 with rhizopine and
341 monitoring activation of the *PnifH::GFP* promoter fusion (S8a Fig). Interestingly, the NifA_{L94/D95Q}
342 protein activated *PnifH::GFP* 13-fold ± [SEM] 1.5 and 98-fold ± [SEM] 2.8 under aerobic and
343 microaerobic conditions, respectively (S8b Fig), suggesting that the protein is tolerant to oxygen.

344



345

346 **Figure 6. Rhizopine control of N_2 fixation, GS adenylation and NH_3 excretion.** (a) Genetic

347 schematic (not to scale) of the low-copy (RK2 replicon) rhizopine receiver plasmid pSIR03. (b) Dose

348 response of GFP induction in *AcLP* ($n = 3$) harbouring pSIR03 with the rhizopine *scyllo*-inosamine

349 (SI) supplemented *in vitro*. Relative fluorescence units (RFU) are defined here as GFP fluorescence /

350 OD_{600nm} . (c) Flow-cytometry analysis of GFP fluorescence in *AcLP* ($n = 4$) harbouring pSIR03

351 incubated for 24-h in the absence or presence of $10 \mu M$ rhizopine. See Table S1 for full statistics. (d)

352 Genetic schematic (not to scale) of the low-copy (RK2 replicon) rhizopine *nifA*_{L94Q/D95Q}-*rpoN*

353 controller plasmid pSIN04. (e) Optimal nitrogenase activity of *AcPU-R22* $\Delta nifA$ carrying pSIN04

354 measured between 5-h – 21-h by acetylene reduction in $n = 6$ cultures grown under microaerobic

355 conditions (3% O_2 in the headspace). (f) Activity of the unadenylylated (active) form of GS in $n = 3$

356 cultures of *AcPU-R22* $\Delta nifA$ carrying pSIN04 incubated for 24-h as determined by γ -glutamyl

357 transferase assays in the presence or absence of 60 mM $MgCl_2$. (g) Spectrophotometric determination

358 of NH_3 in media of $n = 6$ cultures of *AcPU-R22* $\Delta nifA$ carrying pSIN04 grown in N_2 -fixing conditions

359 (N-free UMS media with 3% O_2 in the headspace) in the presence or absence of $10 \mu M$ SI. Error bars

360 represent one SEM. Independent two-tailed students t-tests with the Bonferroni-holm adjustment were

361 used to compare means. Exact P-values are provided where $P > 0.5$. ** $P < 0.01$, *** $P < 0.001$, **** P
362 < 0.0001 .

363

364 We finally performed NH_3 excretion assays on cultures of strain *AcPU-R22* carrying pSIN04
365 supplemented with 10 μM SI and demonstrated that the strain excreted $812.58 \pm [\text{SEM}] 5.59 \text{ uM}$
366 $\text{OD}_{\lambda 600\text{nm}}^{-1} \text{NH}_3$ into the growth media after 24-h incubation in N_2 -fixing conditions at an optimal
367 rate of $65.13 \pm 7.35 \text{ uM OD}_{600\text{nm}}^{-1} \text{h}^{-1}$ (Fig 5g). Importantly, no NH_3 excretion was detected in the
368 absence of SI, demonstrating tight control. Overall, these experiments confirmed that we had
369 established rhizopine control of N_2 -fixation, GS adenylation and NH_3 excretion in our engineered
370 *AcLP* strain.

371

372 **Discussion**

373 In this study, we employed two strategies to interfere with GS and stimulate NH_3 excretion in *AcLP*.
374 For our first strategy, we attempted to recapitulate previous experiments where insertional inactivation
375 of the P_{II} genes *glnB* and *glnK* stimulated shutdown of GS by adenylation and alleviated negative
376 feedback inhibition of nitrogenase by the product NH_3 , preventing NH_3 assimilation and favouring
377 excretion into the growth media (28). Although we could delete either of the *glnB* or *glnK* genes from
378 *AcLP*, we were unable to delete both genes in the same strain unless a second copy of *glnB* was first
379 introduced into the chromosome, suggesting at least one of the P_{II} coding sequences was essential for
380 growth. Considering that a *Paph::KIXX* kanamycin resistance cassette was previously inserted to the
381 3'-end of the *Ac glnB* coding sequence (25) leaving most of the 5'-end reading frame intact, it seems
382 possible that the GlnB protein may have retained some essential function unrelated to AT and GS
383 activity. In contrast, previous insertion of the omega interposon into *Ac glnK* replaced a segment of
384 the internal coding sequence and was therefore more likely to have abolished the function of the
385 protein (53). Interestingly, similar *glnB* and *glnK* antibiotic cassette insertions have been made in the
386 phototrophic diazotroph *Rhodobacter capsulatus*, resulting in NH_3 -insensitive NifA and nitrogenase

387 expression and activity (54). However, attempts to delete both genes were also unsuccessful in this
388 bacterium (55). Regardless of why deleting *glnB* and *glnK* is lethal, reproducing exact copies of the
389 original *glnB* and *glnK* mutants (28) would likely be required to establish control of NH₃ excretion in
390 *AcLP*, as we have shown here that deletion of *glnK* paired with strong repression of *glnB* had minimal
391 effect on nitrogenase or GS regulation and permitted only low-level NH₃ excretion.

392

393 As was previously demonstrated in *A. brasilense* (29), expression of *E. coli* or *Ac*-derived uATs in our
394 *AcLPΔglnE* mutant resulted in strong GS shutdown and high rates of NH₃ excretion when grown in
395 N₂-fixing conditions. We also found that while nitrogenase expression and activity is repressed in
396 microaerobic NH₃ or glutamine-fed cultures of *AcLP*, shutdown of glutamine biosynthesis by uAT
397 expression resulted in nitrogenase expression that was unimpeded by NH₃ but still repressed by
398 glutamine, suggesting that NH₃ must first be converted to glutamine or potentially other amino acids
399 such as asparagine (56) to facilitate repression. This same effect was previously reported in
400 phototrophic *Anabaena* spp (57, 58) and *Rhodobacter sphaeroides* (59) where GS activity was
401 shutdown using the chemical inhibitor L-Methionine sulfoximine, and in *Klebsiella pneumoniae*
402 mutants unable to grow on NH₃ as a sole source of N (60). Moreover, In *R. capsulatus*, where N₂-
403 fixation is repressed in response to added NH₃ at three levels; a) NtrC-dependent transcription of *nifA*;
404 b) NifA-dependent transcription of nitrogenase; and c) DraT-DraG-dependent ADP ribosylation of
405 nitrogenase (61, 62); all three levels of regulation were non-responsive to NH₃ following shutdown of
406 GS by insertional inactivation of both P_{II} genes (54). Thus, it seems plausible that shutdown of
407 glutamine biosynthesis from NH₃ and glutamate abolishes NH₃-dependent regulation of N₂-fixation in
408 genetically diverse bacteria. Targeted GS shutdown therefore affects NH₃ excretion on two fronts,
409 allowing sustained nitrogenase expression and activity in the presence of fixed N₂ and preventing
410 assimilation of NH₃, favouring excretion into the environment.

411

412 Without establishing control of GS shutdown, engineered NH₃ excreting diazotrophs are typically
413 auxotrophic for glutamine, which would render them non-competitive in the environment (13, 16).
414 Here, we placed expression of the *uAT-Ac2* allele under control of the NifA-inducible nitrogenase

415 promoter *PnifH* which, when tuned correctly, triggered GS shutdown and NH₃ excretion specifically
416 under N₂-fixing conditions. In the field, this could allow bacteria to retain competitiveness prior to
417 forming O₂-deplete biofilms on the surface of roots (63), however lack of host-specific control could
418 permit provision of NH₃ to non-target plant species. Thus, we further modified the engineered strain
419 AcPU-R22 by deleting *nifA* and bringing the mutant *nifA*_{L94Q/D95Q} and *rpoN* alleles under rhizopine-
420 inducible control, permitting *in vitro* rhizopine-dependent activation of nitrogenase activity, GS
421 shutdown and NH₃ excretion. The resulting strain should also only activate these processes when
422 colonising the roots or rhizosphere of transgenic *RhiP* barley (47), though we acknowledge that
423 experimental demonstration of this will first require optimisation of the current rhizopine signalling
424 circuitry that permits perception of rhizopine by 10-25% of cells colonising *RhiP* barley roots and
425 activation of nitrogenase activity equating to approximately 15% of that observed for wild-type AcLP
426 cells colonising wild-type barley (46). We have made progress on improving rhizopine signalling here
427 as our new low-copy rhizopine receiver plasmid pSIR03 permitted rhizopine-responsive GFP
428 activation in over 90% of cells in the population, compared to our previously reported plasmid
429 pSIN02 where only 36% of cells perceived rhizopine *in vitro* (46). Moving forward, increasing the
430 proportion of *Ac* cells sensing rhizopine on the root surface and rhizosphere will be crucial to
431 establish effective control of traits.

432

433 With bacteria engaging in partner-specific activation of NH₃ excretion, there still exists the possibility
434 that viability and competitiveness might suffer due to increased energy demand (14). Rhizobia
435 overcome this problem by engaging in stringent signalling with the legume that permits partner-
436 specific infection of nodules (64). Inside the nodule, the bacteria are provided with low-oxygen
437 conditions conducive to nitrogenase stability, they can escape the fierce competition of the
438 rhizosphere, and are fed carbon in the form of dicarboxylates (50, 51). Engineering a nodule-like
439 niche with stringent entry requirements into cereals will likely be important to maximise the
440 effectiveness of inoculation with engineered NH₃ excreting inoculants. The strains developed here
441 could be adapted for entry of such an environment and therefore, this work represents significant
442 advancement towards the development of both associative and more intimate “synthetic N₂-fixing

443 symbiosis” with cereals.

444

445 **Materials and methods**

446 **Bacterial strains and plasmids**

447 Bacteria used in this study (S1 File) were cultured in TY (65) or UMS (66, 67) media supplemented
448 with 300 μ M nicotinic acid and 20 mM succinate as previously described (46). Plasmids (S2 Table)
449 were constructed using HiFi assembly (New England Biolabs) or BEVA modular golden-gate
450 assembly (68, 69) as outlined in the S1 File and were mobilised into *Azorhizobium* by diparental
451 mating with *E. coli* ST18 (70). For mini-Tn7 integration into the chromosome, tri-parental matings
452 were used to additionally mobilise the transposase helper plasmid pTNS3, which carries an R6K
453 origin of replication that is not maintained in *Azorhizobium* (71).

454

455 Gene deletion and replacement mutant strains were constructed by mobilising the relevant suicide
456 plasmid, derived from pK19mobSacB (S2 Table & S1 File), into the target strain and selecting for
457 single-crossover integration into the chromosomal region of interest by plating cells on selective UMS
458 or TY agar media supplemented with 100 μ g mL⁻¹ kanamycin. Single-crossover mutants were
459 subsequently grown in non-selective media until stationary phase and plated in serial dilutions onto
460 UMS or TY agar supplemented with 10% (v/v) sucrose to select for double crossover deletion or
461 replacement of the target gene. For the *Δ glnK:: Ω Sp* replacement plasmid pOPS1564 only, 100 μ g mL⁻¹
462 spectinomycin and 1 mM IPTG was added to the media unless otherwise stated. Single colonies
463 were patched onto the same media used for double-crossover selection plus and minus 100 μ g mL⁻¹
464 kanamycin and kanamycin sensitive colonies were screened by PCR and sanger sequencing for
465 deletion or replacement of the target gene.

466

467 All AcLP *Δ glnB Δ glnK:: Ω Sp* mutant strains were constructed by first deleting *glnB* from AcLP using
468 plasmid pOPS1691, then subsequently integrating the *Δ glnK:: Ω Sp* replacement plasmid pOPS1564

469 into the target chromosomal region by single-crossover. Because replacement of *ΔglnK::ΩSp* was not
470 possible on three separate occasions, mini-Tn7 delivery plasmids carrying an IPTG-derepressible
471 copy of *glnB* (S1 File) were integrated into the engineered *attB* site prior to selecting for selecting for
472 double-crossover replacement of *glnK* with the *ΩSp* interposon as described above.

473

474 **Growth curves**

475 Growth curves were performed in triplicate by streaking single colonies onto 10 mL TY agar slopes
476 and incubating for 3-days prior to three washes in PBS and inoculation at OD_{600nm} 0.01 into 500
477 μL UMS media in 24-well plates. The OD_{600nm} was monitored at 20 min intervals in an Omega
478 FLUOstar plate reader set to shake cultures at 700 rpm at 37 °C until stationary phase. Growth
479 statistics were calculated using the R package GrowthCurver (72).

480

481 **Glutamine synthetase transferase assays**

482 Six-millilitre UMS cultures were initially grown in 30 mL glass universal vials sealed with silicone
483 rubber septa as described for RT-qPCR experiments. After 3-h or 24-h incubation, 1 mL of culture
484 was sampled for protein determination using a Millipore BCA protein assay kit. Five hundred
485 microlitres of CTAB (1 mg mL⁻¹) was added to the remaining cultures which were incubated at room
486 temperature for a further 3 mins prior to harvesting by centrifugation at 4 °C. Cells were washed once
487 with 5 mL 1% (w/v) KCL and finally resuspended in 500 μL of the same buffer and stored on ice. GS
488 transferase assays were performed on 50 μL aliquots the permeabilised cells as previously described
489 (15). The assays were performed in 500 μL total volumes with 30 min incubation in the presence or
490 absence of 60 μM added Mg²⁺ to determine the total GS transferase activity and the activity of the
491 “active” unadenylylated enzyme, respectively (48). The GS transferase buffer was adjusted to pH 7.0,
492 as this was previously estimated as the iso-activity point for *Ac* (73). Following addition of the FeCl₃
493 stop reagent, reaction tubes were centrifuged for 5 min at 13,000 g and 200 μL was transferred to
494 clear, flat bottomed 96-well plates for spectrophotometric quantification of the product L-Glutamyl-γ-
495 Hydroxamate (LGH) at 562nm in a Promega GloMax multi-detection system.

496

497 **Acetylene reduction assays**

498 Cultures for ARAs were prepared and analysed as previously described (46, 74) and 1 mL samples of
499 the headspace atmosphere were analysed using a PerkinElmer Clarus 480 gas chromatograph
500 equipped with a HayeSep® N (80–100 MESH) 584 column at 3-h, 5-h, 21-h, 23-h and 25-h
501 incubation, unless otherwise stated.

502

503 **NH₃ excretion assays**

504 Three-millilitre UMS cultures were initially grown in 30 mL glass universal vials sealed with silicone
505 rubber septa as described for RT-qPCR experiments. OD_{600nm} was recorded and NH₃ was
506 quantified in spent supernatants using the spectrophotometric indophenol assay as previously
507 described (15). A calibration curve was performed for each experiment using freshly made dilutions
508 of NH₃Cl in UMS ranging from 5 µM – 1 mM. Absorbance of indophenol blue was quantified in a
509 Genesys 150 UV visible spectrophotometer (Thermo Scientific) at 652nm after 4-h incubation at
510 room temperature.

511

512 **RT-qPCR**

513 For RT-qPCR experiments, *n*=5 single colonies were streaked onto 10 mL UMS agar slopes
514 supplemented with 20 mM succinate, 10 mM NH₄Cl and 300 µM nicotinate and grown for 2-days at
515 37 °C. Cells were washed three times from the slopes with PBS, resuspended in UMS supplemented
516 with the relevant carbon and N sources at OD_{600nm} 0.3 in 30 mL glass universal vials and
517 transferred with the lid off into a sealed atmosphere cabinet adjusted to 3% O₂ by flushing with N₂
518 gas. After 30 min, cultures were sealed with silicone rubber septa and incubated at 37 °C with
519 rigorous shaking for 3-h. Cells were next harvested by centrifugation at 4 °C, lysed using a FastPrep-
520 24 5G instrument and cellular debris was removed by a second round of centrifugation. RNA was
521 extracted from the resulting lysate using a Qiagen RNAeasy extraction kit and tested for quality and

522 purity using an Agilent Experion Bioanalyzer with RNA Stdsens chips. gDNA was depleted from
523 RNA by treatment with Invitrogen Turbo DNase as per the manufacturer's recommendations and 5
524 µg was used to generate cDNA using an Invitrogen SuperScript IV reverse transcriptase kit as per the
525 manufacturer's recommendations. The final cDNA template was diluted 1:20 with water and 1 µL
526 was added to each 20 µL RT-qPCR reaction prepared in 96-well plates with Applied Biosystems
527 PowerUp SYBR green master mix. Reactions were run using an Applied Biosystems ViiA 7 Real-
528 Time PCR system. RT-qPCR primers were initially tested for amplification efficiency and target
529 specificity by generating a standard curve of amplification with 5-fold dilutions of *AcLP* gDNA. The
530 housekeeping gene primer targeted *recA* and was validated previously (75), whereas the *glnA* primers
531 designed here had the following sequence *glnA F* 5'- CCGCTGACCAACTCCTACA *glnA R* 5'-
532 CCATGAACAGGGCCGAGAA.

533

534 **GFP reporter assays and flow-cytometry**

535 GFP reporter assays and flow-cytometry experiments were performed on 24-h incubated cultures as
536 previously described (46). Inducers were added directly to the growth media at the time of inoculation
537 where relevant.

538

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542

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720 **Supporting Information Captions**

721 **S1 Fig. Characterisation of synthetic ribosome binding sites in *AcLP*.** Each RBS was fused to GFP
722 under expression by the strong synthetic promoter J23104 on plasmid pOGG024 and GFP was
723 measured after 24-h incubation in UMS media ($n = 3$). Relative luminescence units are defined here
724 as GFP fluorescence / OD₆₀₀λnm. The RBS nucleotide sequences are provided in S1 File.

725

726 **S2 Fig. Expression and total activity of glutamine synthetase (*glnA*, GS) is elevated in *AcRGI*.** (a)
727 Total specific activity of both adenylylated (inactive) and unadenylylated (active) forms of GS was
728 measured in whole cells grown for 24-h as determined by γ -glutamyl transferase assays ($n = 5$). (b)
729 *glnA* expression was quantified relative to the housekeeping gene *recA* by RT-qPCR in cells growth
730 for 3-h. All cultures for assays were grown in N₂-fixing conditions (N-free UMS media with 3% O₂ in
731 the headspace). Error bars represent one SEM. Independent two-tailed students t-tests were used to
732 compare means. ***P < 0.001.

733

734 **S3 Fig. Induction of the *Sinorhizobium meliloti* 1021 naringenin-inducible *PnodA* promoter in**
735 ***AcLP*** (a) Genetic schematic (not to scale) of the low-copy (RK2 replicon) naringenin-inducible *GFP*
736 reporter plasmid pOPS1536. (b) GFP induction in *AcLP* ($n = 3$) harbouring pOPSXXX in response to
737 naringenin supplemented *in vitro*. Relative luminescence units are defined here as GFP fluorescence /
738 OD₆₀₀λnm.

739

740 **S4 Fig. Total activity of glutamine synthetase (GS) in Δ *glnE* mutants expressing unidirectional**
741 **adenylyl transferases from the non-induced *PnodA* promoter** (a) Total specific activity of both
742 adenylylated (inactive) and unadenylylated (active) forms of GS was measured in whole cells grown in

743 N₂-fixing conditions (N-free UMS media with 3% O₂ in the headspace) for 3-h as determined by γ -
744 glutamyl transferase assays ($n = 5$ for wild-type *AcLP* or $n = 3$ for other strains). Error bars represent
745 one SEM. Independent two-tailed students t-tests were used to compare means against the wild-type
746 (WT) *AcLP* as a reference. Not significant (ns) indicates $P > 0.05$, * $P < 0.05$.

747

748 **S5 Fig. NH₃ excretion is sub-optimal in $\Delta glnE$ mutants expressing unidirectional adenylyl**
749 **transferases from the *PnodA* promoter induced with naringenin.** Spectrophotometric
750 determination of NH₃ in media of cultures induced with 5 μ M naringenin grown for 24-h in N₂-fixing
751 conditions (N-free UMS media with 3% O₂ in the headspace). Error bars represent one SEM. $n = 3$
752 for wild-type *AcLP* $\Delta glnE$ or $n = 6$ for other strains.

753

754 **S6 Fig. Growth statistics for control strains and $\Delta glnE$ mutants expressing unidirectional**
755 **adenylyltransferases.** Mean generation times and the max OD_{600nm} (i.e. the carrying capacity, k)
756 were calculated from standard curves of cultures grown in UMS media at 21% O₂. Representative
757 growth curves are provided in Fig 3c and Fig 5b. Strains highlighted in white are wild-type (WT)
758 *AcLP* and *AcLP* $\Delta glnE$ controls, strains highlighted in pink are *AcLP* $\Delta glnE$ carrying *PnodA* [RBS]
759 uAT-DT16 modules on parent plasmid pOGG093 and strains highlighted in blue are *AcLP* $\Delta glnE$
760 carrying mini-Tn7 integrated *PnifH* [RBS] uAT-*Ac2*-DT16 modules.

761

762 **S7 Fig. Rhizopine control of N₂ fixation alone does not permit NH₃ excretion.** Spectrophotometric
763 determination of NH₃ in media of $n = 3$ cultures grown for 24-h in N₂-fixing conditions. Error bars
764 represent one SEM. Strain *Azospirillum brasilense* HM053 was used here as a positive control.

765

766 **S8 Fig. Nif_{L94QD95Q} activity is tolerant to ambient environmental oxygen. (a)** Genetic schematic
767 (not to scale) of the rhizopine *nif_{L94QD95Q}-rpoN* controller plasmid with *PnifH::GFP* reporter fusion
768 pSIN03. **(b)** *PnifH* promoter activity was measured in $n = 4$ cultures grown for 24-h under the
769 conditions indicated. Relative fluorescence units (RFU) are defined here as GFP fluorescence /

770 OD600λnm. Error bars represent one SEM. Independent two-tailed students t-tests with Bonferroni-
771 holm adjustment were used to compare means. $P > 0.05$. ** $P < 0.01$, *** $P < 0.001$.

772

773 **S1 Table. Flowcytometry statistics for rhizopine-inducible GFP expression in *AcLP* carrying**
774 **pOPS1052**

775

776 **S2 Table. Plasmids used in this study**

777